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## Mutations in outer dynein arm heavy chain *DNAH9* cause motile cilia defects and situs inversus

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## Abstract

Motile cilia move body fluids and gametes and the beating of cilia lining the airway epithelial surfaces ensures that they are kept clear and protected from inhaled pathogens and consequent respiratory infections. Dynein motor proteins provide mechanical force for cilia beating. Dynein mutations are a common cause of primary ciliary dyskinesia (PCD), an inherited condition characterised by deficient mucociliary clearance and chronic respiratory disease coupled with laterality disturbances and subfertility. Using next-generation sequencing we detected mutations in the ciliary outer dynein arm (ODA) heavy chain gene *DNAH9* in individuals from PCD clinics with situs inversus and in one case male infertility. *DNAH9* and its partner heavy chain *DNAH5* localise to type 2 ODAs of the distal cilium and in *DNAH9*-mutated nasal respiratory epithelial cilia we found a loss of *DNAH9/DNAH5*-containing type 2 ODAs that was restricted to the distal cilia region. This confers a reduced beating frequency with a subtle beating pattern defect affecting the motility of the distal cilia portion. 3D electron tomography ultrastructural studies confirmed regional loss of ODAs from the distal cilium, manifesting as either loss of whole ODA or partial loss of ODA volume. *Paramecium DNAH9*-knockdown confirms an evolutionarily conserved function for *DNAH9* in cilia motility and ODA stability. We find *DNAH9* is widely expressed in the airways, despite *DNAH9* mutations appearing to confer symptoms restricted to the upper respiratory tract. In summary, *DNAH9* mutations reduce cilia function but some respiratory mucociliary clearance potential may be retained, widening the PCD disease spectrum.

Cilia are highly conserved microtubular organelles found on almost every cell of the human body.{

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 cilia line the upper and lower respiratory tract, ependymal cells, Fallopian tubes and the Eustachian tubes.  
 Singleton motile cilia present during early development function at the embryonic left-right organiser,  
 also known as the node. These different motile cilia beat in a co-ordinated fashion to transport overlying  
 fluid. The single flagellum of sperm tails is similar in structure to motile cilia.{ ADDIN EN.CITE {  
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Primary ciliary dyskinesia (PCD, MIM#244400) is an autosomal recessive or X-linked condition in which  
 the motile cilia are dysfunctional and fail to effectively transport secretions.<sup>3; 4</sup> Pathogenic variants in >  
 35 genes involved in cilia assembly, structure and motility have been identified to cause PCD.<sup>5; 6</sup> In  
 keeping with the function and location of motile cilia and sperm, symptoms of PCD can include neonatal  
 respiratory distress, chronic productive cough, recurrent infections of the upper and lower airways,  
 middle ear symptoms such as otitis media and male and female infertility. Due to cilia dysfunction in the  
 embryonic node, situs inversus and other congenital cardiac and laterality disorders also manifest.<sup>3; 6</sup>  
 Diagnosis typically uses a combination of clinical tests including nasal nitric oxide (nNO) measurement,  
 ciliary beat and ultrastructure analysis by high speed video and transmission electron microscopy  
 (HVSM, TEM), immunofluorescence staining of motile cilia proteins and genetic analysis.<sup>6</sup>

The core structure (axoneme) of motile cilia consists of a circle of nine peripheral microtubular doublets  
 plus or minus a central pair of microtubules, respectively termed ‘9+2’ (multiple cilia) or ‘9+0’ (singleton  
 nodal cilia).<sup>1; 4</sup> Inner (IDA) and outer dynein arms (ODA) that drive ciliary beating through ATP

hydrolysis are attached to the peripheral microtubular doublets. The mammalian ODA, a complex structure attached to the microtubules by a docking complex, is composed of multiple heavy, light and intermediate dynein chains.<sup>7; 8</sup> In a highly evolutionarily conserved fashion, the ODA structure repeats every 24nm along the length of the axoneme and is controlled through the nexin dynein regulatory complex and radial spokes to produce a regular ciliary waveform.<sup>9; 10</sup> Two types of human ODAs are described.<sup>11</sup> Type 1 ODA reside in the proximal part of the cilium closer to the cell body in the region of the microvilli, containing the two dynein heavy chains (HC) DNAH5 (orthologous to the ODA  $\gamma$ -HC found in *Chlamydomonas*, the ancient protist that forms a key PCD model system) and DNAH11 (orthologous to the *Chlamydomonas* ODA  $\beta$ -HC). Type 2 ODA reside in the distal cilium beyond the region of the microvilli, containing the two heavy chains DNAH5 and DNAH9 (also orthologous to *Chlamydomonas* ODA  $\beta$ -HC).<sup>11; 12</sup>

Mutations in *DNAH5* (MIM#603335) and *DNAH11* (MIM#603339) are amongst the commonest causes of PCD.<sup>13-15</sup> *DNAH5* mutations result in loss of the  $\gamma$ -HC DNAH5 and the  $\beta$ -HC DNAH9 and consequently a classic ODA loss visible by TEM. In contrast, pathogenic variants in *DNAH11* result in apparent normal ultrastructure by TEM, with ODAs continuing to assemble DNAH5 and DNAH9 despite the absence of DNAH11.<sup>14</sup> Higher resolution ultrastructural analysis using electron tomography reveals that *DNAH11* mutations do confer a subtle structural defect, affecting only the proximal type 1 ODAs with a reduced ODA volume.<sup>16</sup> Furthermore, in individuals with pathogenic variants in *DNAH11*, DNAH9 can reside in the place of DNAH11 in the proximal part of the axoneme.<sup>12</sup>

Here, we describe individuals with mutations affecting highly conserved domains of the ODA heavy chain *DNAH9* (MIM#603330). We report clinical, genetic, immunofluorescence, ciliary beat analysis and electron tomography ultrastructural studies that increase our understanding of the arrangement of

type 2 ODA heavy chains of motile cilia. These reveal that *DNAH9* mutations confer a loss of ODAs in the distal portion of the ciliary axoneme, accompanied by a subtle effect on the ciliary beat pattern.

This study was approved by the ethics review boards of the Institute of Child Health/Great Ormond Street Hospital, London (UK) (08/H0713/82) and the Comité de Protection des Personnes CPP Ile-de-France III (France) (CPP07729 and CPP02748). We used a targeted next generation sequencing (NGS) gene panel (Agilent SureSelectQXT or Roche SeqCap EZ Choice) for genetic screening in a total of 536 individuals with confirmed or suggestive diagnosis of PCD. We identified loss-of-function mutations in *DNAH9* (GenBank: NM\_001372.3) in four individuals from three unrelated families (**Figure 1A**). Family 1 has one male child born in the UK to Turkish consanguineous parents with rhinosinusitis, cough, situs inversus and complex congenital heart disease consisting of an unbalanced atrioventricular septal defect with congenitally corrected transposition of the great arteries and inferior vena cava conduit stenosis that required stenting. He also has a gastrointestinal condition (protein-losing enteropathy). He carries a homozygous missense mutation c.12367G>A; p.(Asp4123Asn), confirmed by Sanger sequencing which also showed the father was a carrier (mother's sample not available) (**Figure 1A**). This variant is absent from the Genome Aggregation Database (gnomAD). Phylogenetic analysis showed that Asp4123 is an amino acid highly conserved across different species and also across several dynein heavy chains. (**Figure 1B**) This variant is predicted to be probably damaging using Polyphen-2 with a score of 1.0 and it has a Combined Annotation Dependent Depletion (CADD) score of 33 (CADD score of  $\geq 20$  indicates a variant is among the top 1% of damaging variants in the genome).

In family 2, a UK based non-consanguineous Somali family with an affected brother and sister both of whom had rhinosinusitis and situs inversus, we found compound heterozygous variants carried by both affected siblings: a missense variant c.10193G>T; p.(Arg3398Leu) and an exon 46 consensus splice



acceptor site variant c.8708-2A>G. Two unaffected siblings carried only the c.10193G>T; p.(Arg3398Leu) allele in a heterozygous state, results confirmed by Sanger sequencing in all four siblings (**Figure 1A**). It is absent from the gnomAD, dbSNP and EVS control databases and predicted by Polyphen-2 to be probably damaging with a score of 0.997 and a CADD score of 34. Arg3398 is a highly conserved amino acid in different species and also across several dynein heavy chains (**Figure 1B**). The c.8708-2A>G splice variant is in dbSNP (rs143007518) and predicted to cause a frameshift, with a CADD score of 23.3. It was confirmed by Sanger sequencing in the two affected siblings and found to be absent in two unaffected siblings. It is present once in the gnomAD control database in a heterozygous state in a carrier from the East Asian population with a total allele frequency of 3.234e-5. To assess its functional impact on *DNAH9* splicing, we isolated total RNA using Trizol Reagent (QIAGEN) from non-cultured ciliated cells obtained through nasal brushing of one of the affected siblings (2.II.3) and a healthy control. We used the High-Capacity RNA-to-cDNA™ Kit (ThermoFisher Scientific) to synthesize cDNA. Using primers in exons 41 and 47 of *DNAH9*, RT-PCR yielded one band of 1,032 basepairs (bp) in the healthy control while two bands (1,032 and 889 bp) were amplified in the affected child. We Sanger sequenced the two bands amplified in the latter sample after purification using the Monarch® DNA Gel Extraction Kit (New England BioLabs). Sequence analysis revealed skipping of exon 46 leading to a predicted shifting of the expressed protein reading frame, producing a stop codon 53 residues downstream [p.(Glu2904Aspfs\*53)] (**Figure 1C**).

In family 3, the Algerian male proband born to a consanguineous union had rhinosinusitis, cough, situs inversus and marked asthenozoospermia (sperm immotility) in fertility testing (**Table S1**). He was identified to carry two homozygous missense mutations, which were confirmed by Sanger sequencing in the affected individual's DNA (**Figure 1A**). No DNA was available from other family members but despite the lack of parental samples, intra-run comparative analysis of the subject's NGS read depth ratio

confirmed that two copies of all the *DNAH9* coding exons were present in his genome, showing that there was no hemizygosity and that is a complex allele with two homozygous variants present (**Figure S1**). The first homozygous missense variant c.5641A>G; p.(Lys1881Glu) affects a highly conserved amino acid flanking the only ATPase site thought to retain an ATP hydrolyse activity (AAA1 domain) (**Figure 1B**) and it is absent from dbSNP, gnomAD and EVS control databases. The CADD score is 27.8 and Polyphen-2 predicts it to be probably damaging with a score of 1.0. The second homozygous missense variant c.8894G>A; p.(Arg2965His) was found in 16 unaffected carriers in gnomAD with a total allele frequency of 5.8e-5 (rs375908701). It changes a highly conserved residue Arg2965 in DNAH9 and this is predicted to be probably damaging with a score of 0.99 in Polyphen-2 and a CADD score of 34 (**Figure 1A, B**).

**Table 1** shows clinical and PCD test results from these individuals, revealing a comparatively mild phenotype. Notably, all affected individuals had situs inversus. Although showing upper respiratory tract symptoms (rhinitis), none of the affected children had recorded signs of neonatal respiratory distress, otitis media or signs of permanent lung damage (bronchiectasis). Their nasal nitric oxide levels were within or close to the normal range (>77nl/min). Three of the four affected probands had lung function (forced expiratory volume in one second (FEV1) % predicted) in keeping with that predicted in healthy children based on their age, height, gender and ethnicity. <sup>6</sup> Individual 1.II.1 had compromised lung function (49% predicted), which is likely attributed to co-morbidities and poor technique in conducting the test.

Two or more nasal brush biopsies per individual were analysed on separate occasions to assess ciliary function. Ciliary beat frequency, measured by HSVM, was at the lower end of the normal range<sup>17</sup> in all individuals (**Table 1**). As summarised in **Figure 2A** and **Figure S2**, ciliary beat pattern analysis revealed

cilia with a subtle defect in the bend of the cilia in the distal portion captured in affected individuals 1.II.1, 2.II.2 and 3.II.1 (**Videos S1-S3**), compared to healthy controls (**Video S4**). Quantitative re-analysis of videos of ciliary beating in nasal cells from affected individuals at one centre (B.L.) confirmed a lower beat frequency associated with the altered beat, and potentially with a lower distance and area swept by the cilium (**Table S2**). As a measure of mucociliary clearance, the transport of latex microbeads across the cilia surface was also assessed on a fresh nasal sample from 3.II.1 compared to a healthy control (**Videos S5 and S6**). The ciliated edges of 3.II.1 clearly induced fluid flow, observed by displacement of the microbeads calculated as a ciliary beating efficiency index <sup>18</sup> of  $0.35 \pm 0.16$  mPa (evaluated on four ciliated edges), lower than the control ( $2.6 \pm 1.3$  mPa, evaluated on 11 edges). This implies some respiratory mucociliary clearance potential may be retained by the cilia in individuals with *DNAH9* mutations. Interestingly, in contrast to their respiratory cilia motility, sperm analysis performed in individual 3.II.1 showed oligozoospermia with only 3% sperm having preserved motility (other affected individuals were too young to be assessed).

We next performed cilia immunofluorescence staining to investigate the effect of *DNAH9* mutations on *DNAH9* localisation in nasal epithelial cilia. In controls, as expected, *DNAH9* was restricted to the distal portion of the cilia (**Figure 2B**). We found similar results in unaffected siblings of family 2 (data not shown). In contrast, affected individuals carrying *DNAH9* mutations showed heavily reduced / absent staining of *DNAH9* protein in the cilia, confirming the severity of the inherited mutations (**Figure 2A**). Since affected individuals in all three families carried single amino acid substitution ‘missense’ changes but showed this marked reduction in ciliary *DNAH9* protein, we performed quantitative RT-PCR to investigate the abundance of *DNAH9* transcript levels. For two cases where nasal sample was available, 2.II.3 and 3.II.1, we found that *DNAH9* transcripts were still present but at variable levels presumably reflecting variable levels of ciliation in primary nasal samples as well as possible mutational effects

(**Figure 1C, Figure S3**). Western blot of the 3.II.1 sample also confirmed a complete lack of the DNAH9 protein (**Figure S3**).

We further sought to identify the location of other ODA components. In controls, DNAH5 stains the entire cilia length (**Figure 2C**) but in cilia from individuals with *DNAH9* mutations, type 2 ODAs in the distal portion of the mutant cilia were lacking DNAH5 staining; this was severely reduced and usually completely absent (**Figure 2C**). DNAH5 staining was unaffected, being similar to controls, in type 1 ODAs in the proximal portion of the cilium (**Figure 2C**). The difference in proximal and distal DNAH5 signal along the length of the cilia from an affected individual with *DNAH9* mutations could be quantified by plotting pixel intensity across 10 profile plots of immunofluorescent antibody labelled cilia (**Figure 2D**). These findings suggest DNAH9 absence often results in loss or failure of assembly of DNAH5  $\gamma$ -HC in the distal part of the cilia.

We analysed the localization of other key motile ciliary components in individuals with *DNAH9* mutations. We found that staining of the ODA intermediate chain DNAI1<sup>8</sup> was altered in affected individuals from all three families carrying *DNAH9* mutations, being lost from the distal part of the cilia only, in parallel to the loss of DNAH9 and DNAH5 (**Figure 2B, C and Figure S4**). Quantification of immunofluorescent antibody pixel intensities along the cilia illustrate these results for the cilia of individual 2.II.3 compared to a control (**Figure S5**). In individual 2.II.2 with *DNAH9* mutations, DNAH11, the other axonemal  $\beta$ -HC present in type 1 ODAs, was present in the proximal portion of the cilium according to its usual localisation behaviour<sup>12; 16</sup> and in a similar distribution compared to controls, as judged by immunostaining (**Figure S6** and quantified in **Figure S5**). This suggests that DNAH11 does not replace DNAH9 in the type 2 ODAs in the absence of DNAH9, however further quantitative work is required to prove this. In contrast CCDC114, a component of the ODA docking

complex, was still present as normal throughout the length of the ODA, as confirmed also by Western blot, suggesting there is no failure in docking complex assembly (**Figure S3 and Figure S7**). Key components of the inner dynein arm and radial spoke head were also present as normal throughout the axoneme as measured by antibodies to DNALI1 and RSPH4A (**Figure S7, Figure S8** and quantified in **Figure S5**).

We next assessed by TEM the ultrastructural effect of *DNAH9* mutations on cilia. Overall, evaluating >100 cilia cross section without bias towards measurements in any particular cilia region, ODA defects affected 35-89% of axonemes in individuals carrying *DNAH9* mutations with significant numbers of ODAs in most cases remaining undisturbed (**Figure 3A**). We then divided the TEM cross sections into those taken from either the distal or the proximal cilia regions. The proximal axonemal cilia segment corresponding to the microvilli region (the location of type 1 ODAs) were of normal composition, whilst axonemes in the distal part of the cilium (judged by distance from the cell surface and absence of surrounding microvilli) had significant numbers of absent or truncated ODAs (**Figure 3B**).

To better understand the nature of *DNAH9*-associated structural defects we performed higher resolution imaging by 3D electron tomography, on axonemes from affected individuals. Tomograms were taken from ODAs that appeared complete by conventional TEM (such as those indicated by black arrows in **Figure 3B**). This confirmed the regionality of the *DNAH9* mutational effect since in individual 1.II.1, tomography revealed a loss of volume only of the type 2 ODA i.e. only ODA in the distal cilium, whilst in the proximal cilium the ODA volume was equivalent to that of controls. For type 2 ODA of the distal cilium that were not completely lost but retained, the reduction of ODA volume was specific to their ‘forearm’ portion, reflecting the predicted location of the *DNAH9*  $\beta$ -HC within the ODA (**Figure 3C**). The ODA ‘forearm’-specific defect was also confirmed by tomography of the ciliary distal region ODAs

in individuals 2.II.2 and 2.II.3 (**Figure S9**). This DNAH9 data is in keeping with our previously published converse (proximal) findings in individuals with *DNAH11* variants, who have a similarly reduced ODA volume for the DNAH11  $\beta$ -HC but affecting only the type 1 ODA of the proximal cilium.<sup>16</sup> Quantification of the ODA volumes in cilia from three *DNAH9*-mutated individuals was compared to results from individuals without respiratory disease. This showed a significant decrease in the ODA volume in the distal portion of their cilia, while ODA volume in the proximal portion of the cilia was not affected (**Figure 3D**). Preliminary tomography studies in individuals with PCD carrying *DNAH5* mutations show some evidence that *DNAH9* mutations may confer a less reduced type 2 ODA volume than *DNAH5* mutations, but further study is required (A.S. unpublished observation). **Figure 3E** presents a model of the findings arising from the current study, showing the regional (distal cilium) effect of DNAH9 loss on the ODA structures in individuals with pathogenic variants in *DNAH9*.

Due to the findings in individuals harbouring *DNAH9* mutations, of upper respiratory tract symptoms but preserved lung function in three of 4 probands and no evidence of bronchiectasis (**Table 1**), we sought to confirm whether DNAH9 was present throughout the respiratory tract and not isolated to nasal tissue. Tissue obtained from an individual without PCD undergoing a lobectomy showed similar localisation of DNAH9, DNAH5 and DNAH11 in the nasal passage, lobar region and the peripheral airways of <1.5mm. This confirms the location of both type 1 and type 2 ODAs in cilia throughout the airways (**Figure 4**).

We recently reported the use of *Paramecium* as a model for motile ciliary disorders, due to the highly conserved nature of ciliary proteins.<sup>19</sup> To study the role of DNAH9 in cilia motility, we employed gene knock-down in *Paramecium* using RNAi silencing by feeding, as described before.<sup>19</sup> We identified two orthologues of *DNAH9* due to whole genome duplication in the *Paramecium* genome (GenBank XM\_001443078.1) (**Figure S10**). Real-time quantitative PCR confirmed >80% knock down of transcript

levels for both these *DNAH9* orthologues by RNAi, comparing *DNAH9*-RNAi *Paramecia* (double knockdown of both orthologues) to *ND7*-RNAi *Paramecia* as a control (the *ND7* gene is routinely used as a control as it has no role in cilia motility<sup>19</sup>) (**Figure 5A**). *Paramecium DNAH9*-RNAi experiments were repeated >3 times to confirm the reproducibility of the results. We showed a significant reduction of the swimming velocity and cilia beating frequency in the *DNAH9* knockdown cells compared to controls over 3 days of RNAi silencing (**Figure 5B-D and Videos S7 and S8**). Ultrastructural loss of ODAs from the axoneme was quantified by TEM (**Figure 5E, F**). The phenotype of the *DNAH9* mutant *Paramecium* cilia therefore recapitulates the human phenotype.

In summary, we report four individuals with PCD carrying mutations in *DNAH9* that affect amino acids in highly conserved protein domains or cause a frameshift, all of which result in severely reduced/absent levels of protein. These individuals all manifest with respiratory symptoms, laterality (situs) defects and male infertility. Our findings are in agreement with Loges et al 2018, who also report mutations in *DNAH9* that cause similar phenotypes.<sup>20</sup> *DNAH9*-related disease is a form of primary ciliary dyskinesia, since the motile cilia of the respiratory system and embryonic node are defective (the latter since situs defects are present<sup>1; 4; 21</sup>) as well as sperm motility.

*DNAH9* is the  $\beta$ -HC dynein of type 2 ODA, which is expressed in the airways and sperm.<sup>11; 22</sup> We find that *DNAH9* is expressed in cilia from all the human upper and lower respiratory airway epithelia we tested. *DNAH9* knock-down in *Paramecium* mimics the ODA and cilia motility defects of *DNAH9*-deficient individuals, indicating the highly conserved critical role *DNAH9* plays in forming the axonemal dynein arms. Recently, *DNAH9* polymorphisms were associated with asthma and bronchial hyperresponsiveness in response to early life tobacco smoke exposure in large GWAS studies, suggesting wider roles in lung pathology.<sup>23</sup>

Our protein localisation and structural studies confirm the previously reported region-specific localisation of DNAH9 in cilia, limited to the distal axoneme portion.<sup>11</sup> In all the affected individuals with PCD, axonemal DNAH9 levels appear heavily reduced to undetectable in immunostaining of their ciliated nasal epithelial cells, even in affected individuals from two families carrying different single amino acid substitution (missense) changes. qRT-PCR showing that *DNAH9* transcripts could still be amplified in these samples, suggest that the mutant forms of the encoded protein may be expressed but not assembled to the axoneme or subject to degradation. This has been noted before for missense mutations causing PCD found in other genes.<sup>24</sup> DNAH9 protein expression was also confirmed to be undetectable by Western blotting of ciliated nasal epithelial cells from 3.II.1 who is homozygous for missense mutations. This suggests these affected protein residues are vital to DNAH9 axonemal localisation and function, supported by their highly conserved nature across other dyneins and DNAH9 orthologues.

The absence of ciliary DNAH9 correlates to a clear loss of type 2 ODA from the airway cilia of individuals carrying *DNAH9* mutations. The heavy and intermediate ODA chains DNAH9, DNAH5 and DNAI1 are all absent/severely reduced from the distal cilia portion, contributing to the regional ODA volume reduction seen by electron tomography. Markers of the axonemal ODA docking complexes, inner dynein arms and radial spokes seem to be unaffected and in place as normal, suggesting the defect is restricted to affecting only the ODAs. *DNAH5* mutations causing PCD have similarly been reported to disturb the ODAs but not the ODA docking complexes.<sup>25</sup>

Type 2 ODA are only located within the distal cilium, such that *DNAH9* mutations are accompanied by a subtle effect on the ciliary beat pattern. We find that individuals carrying *DNAH9* mutations usually have nasal nitric oxide levels in the normal range along with other features of a sometimes mild clinical



presentation. These findings challenge the current paradigm of the phenotype of PCD. *DNAH9* was previously considered as a PCD candidate gene by Bartoloni et al. however no pathogenic mutations were found in 31 PCD families screened before the advent of next generation sequencing.<sup>26</sup> We conclude that *DNAH9* cases can be hard to recognise without genetic testing. Notably, all the individuals carrying *DNAH9* mutations in this study displayed laterality defects (expected to affect 50% of PCD cases) and this may have been a lead symptom for their referral to the PCD clinic.

Interestingly, infertility tests in one available case in our study show that *DNAH9* defects may have a greater impact on sperm motility than cilia motility, however further functional analysis in a larger cohort would be needed to confirm whether this is the case, including further studies of the distribution of *DNAH9* in human sperm compared to cilia.<sup>11</sup> Altogether, these clinical findings can better inform disease diagnostics, disease management and appropriate counselling of affected families.

Previous work showed that in individuals with PCD caused by *DNAH11* defects, whilst *DNAH11* is absent from their cilia, *DNAH9* shifts from a distal to pan-axonemal localization i.e. is aberrantly present in place of the type 1 ODA; however despite this, the shift in *DNAH9* localisation is not able to rescue the *DNAH11* motility defects.<sup>12</sup> In contrast to this pattern, our present study demonstrates no evidence in subjects with *DNAH9* mutations for a change in their ciliary *DNAH11* localisation or its substituting for *DNAH9*, implying that *DNAH11* and *DNAH9*  $\beta$ -HCs behave differently. Interestingly, a recent study proposed a sequential assembly of *DNAH5* and *DNAH9*, followed by *DNAH11* throughout different stages of ciliogenesis towards generation of a mature cilium, so the protein distribution (and ciliary waveform) varies at different stages.<sup>27</sup> In our study, we ensured to select only full-length cilia for quantification of signal intensity of the different ODA markers from the proximal to distal regions.

In conclusion, this study sheds new light on ODA HC biology and the effect of inherited mutations in ciliary ODA genes, broadening our understanding of the PCD clinical phenotypic spectrum.

## Supplemental data

10 figures

5 tables

8 videos

Video S1 – 1.II.1 HSVM nasal cilia

Video S2 – 2.II.2 HSVM nasal cilia

Video S3 – 3.II.1 HSVM nasal cilia

Video S4 – Healthy HSVM control nasal cilia

Video S5 – 3.II.1 microbead transport

Video S6 – Healthy control microbead transport

Video S7 – HSVM *ND7*-knockdown control *Paramecium* cilia

Video S8 – HSVM *DNAH9*-knockdown *Paramecium* cilia

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## Declaration of interests

The authors declare no competing interests.

## Web Resources

Genbank	<a href="https://www.ncbi.nlm.nih.gov/genbank/">https://www.ncbi.nlm.nih.gov/genbank/</a>
GnomAD	<a href="http://gnomad.broadinstitute.org/">http://gnomad.broadinstitute.org/</a>
dbSNP Build 141	<a href="https://www.ncbi.nlm.nih.gov/projects/SNP/">https://www.ncbi.nlm.nih.gov/projects/SNP/</a>
Exome Variant Server	<a href="http://evs.gs.washington.edu/EVS/">http://evs.gs.washington.edu/EVS/</a>
Polyphen-2	<a href="http://genetics.bwh.harvard.edu/pph2/">http://genetics.bwh.harvard.edu/pph2/</a>
Combined Annotation Dependent Depletion	<a href="http://cadd.gs.washington.edu/">http://cadd.gs.washington.edu/</a>
CILDB	<a href="http://cildb.cgm.cnrs-gif.fr/">http://cildb.cgm.cnrs-gif.fr/</a>
ParameciumDB	<a href="http://paramecium.cgm.cnrs-gif.fr/">http://paramecium.cgm.cnrs-gif.fr/</a>
RNAi off-target tool	<a href="http://paramecium.cgm.cnrs-gif.fr/cgi/tool/alignment/off_target.cgi">http://paramecium.cgm.cnrs-</a> <a href="http://paramecium.cgm.cnrs-gif.fr/cgi/tool/alignment/off_target.cgi">gif.fr/cgi/tool/alignment/off_target.cgi</a>
OMIM	<a href="http://www.omim.org/">http://www.omim.org/</a>
RefSeq	<a href="http://www.ncbi.nlm.nih.gov/RefSeq">http://www.ncbi.nlm.nih.gov/RefSeq</a>

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## Figure titles and legends

### Figure 1. Loss of function biallelic *DNAH9* mutations identified in 3 unrelated families

(A) Pedigrees of three unrelated families with *DNAH9* mutations. Family 1 has one affected child harbouring homozygous c.12367G>A; p.(Asp4123Asn) mutations, Family 2 has two affected siblings with compound heterozygous splice acceptor and missense mutations c.8708-2A>G; p.(Glu2904Aspfs\*53) and c.10193G>T; p.(Arg3398Leu), Family 3 has one affected individual with two homozygous missense mutations c.5641A>G; p.(Lys1881Glu), c.8894G>A; p.(Arg2965His). (B) The three identified missense mutations all affect highly evolutionarily conserved residues which are also conserved across most axonemal dyneins. Notably in 3.II.1 carrying two homozygous missense variants, the p.(Lys1881Glu) mutation targets a highly conserved amino acid flanking the only ATPase site thought to retain an ATP hydrolyse activity (AAA1 domain). In addition, amino acid Arg2965 also shows cross species conservation. (C) RT-PCR of nasal epithelial brush biopsy samples yielded one band of 1,032 bp in the healthy control and two bands (1,032 and 889 bp) in the sample from individual 2.II.3 carrying the *DNAH9* c.8708-2A>G putative canonical splice site mutation. cDNA sequencing showed that c.8708-2A>G causes excision of *DNAH9* exon 46 from the transcript and a frameshift mutation: p.(Glu2904Aspfs\*53).

### Figure 2. Loss of distal ciliary ODA type 2 heavy chains (*DNAH9* and *DNAH5*) from the respiratory cilia of individuals with *DNAH9* mutations

(A) Diagram of cilia beating in a control compared to a *DNAH9* mutation case (2.II.2), showing the subtle defect in the bend of the distal portion of the cilia that is captured in HSVM (effective stroke shown in blue, recovery stroke in green). *DNAH9* deficient cilia have reduced beating frequency accompanied by reduced amplitude of the distal cilia area. (B, C) Respiratory epithelial cells were double-stained with an

antibody marker of the ciliary axoneme (acetylated alpha tubulin, green) and the ODA  $\beta$ -HC and  $\gamma$ -HC antibody markers (DNAH9 and DNAH5, red). Co-localisation of acetylated tubulin and presence of dynein heavy chain appears yellow in merged images in the right column. Merged images of the red and green are overlaid with a light microscopy image to aid visualisation of the cell and an image of the nucleus labelled with DAPI in blue. (B) DNAH9 localizes distally in the cilia of control but is absent from cilia of affected individuals. (C) In contrast to the control cilia where DNAH5 localizes along the whole length of the cilia, it is highly reduced in the distal part of the cilia of affected *DNAH9*-mutation carrying individuals but is still present proximally. All scale bars, 5  $\mu$ M. (D) A graph showing DNAH5 signal quantified by plotting the pixel intensity in images across profile plots of immunofluorescent images of antibody labelled cilia, in individual 2.II.3. DNAH5 is present mainly in the proximal part of the cilia and reduced in the distal part (results from control are shown in **Figure S4**)

### **Figure 3. Ultrastructural defects of the distal outer dynein arm in individuals with *DNAH9* mutations**

(A) Brushings of the nasal epithelium were assessed by transmission electron microscopy in four individuals with *DNAH9* mutations and one unaffected sibling (2.II.1). A minimum of 100 axonemes in cross section were assessed per individual. The number of axonemes demonstrating defects of the dynein arms are expressed as a percentage. Both dynein arms were present in the majority of the axonemes assessed (11-65%), the remaining had absent outer dynein arms (89-35%). Where ODAs were absent, usually the accompanying inner dynein arms remained present (orange) but in occasional axonemes the IDA was also absent (grey). The unaffected sibling has 99% normal ultrastructure in keeping with quantification in previously described healthy controls.<sup>28</sup> (B) Example of transmission electron micrographs of ciliary axonemes in cross section from a healthy control subject (1st column) and four individuals with *DNAH9* mutations (columns 2-5). ODAs are highlighted by black arrowheads

in the control and ODA loss is highlighted by the white arrowheads. ODAs are intact in ciliary cross sections judged to be proximal to the cell surface (bottom row) whereas they are frequently absent in the distal part of the cilium (top row). Scale bar = 100nm. (C) Electron tomography and averaging were used to generate 3D models of the ODA, which have been falsely coloured blue for visualisation. Using PEET analysis, transmission electron microscope tomograms taken from an area with >6 ciliary cross sections (54 microtubular doublets) were combined into a single model. Models from tomograms from individual 1.II.1 in the proximal cilia region, judged by the presence of surrounding microvilli, are similar to the models of both the proximal and the distal region of the healthy control axonemes. The model from the distal region of the axonemes of 1.II.1 is reduced in volume (indicated by the black arrow). This model was created from ODAs which appeared to remain intact in standard electron micrographs. (D) The volume occupied by the ODA as a proportion of the entire microtubular doublet distal model was quantified for 3 affected individuals (1.II.1, 2.II.1, 2.II.2) and 6 healthy controls. Data is plotted as box and whisker plots, showing a difference in volume in the distal region compared to controls ( $p > 0.001$ , independent samples t test). Additional tomograms from individuals carrying *DNAH9* mutations used for quantification are shown in the supplementary data (Fig S5). (E) Information from immunofluorescence protein location studies and electron tomography ultrastructural studies were used to create a model of type 1 and type 2 ODAs in cilia from normal controls and individuals with *DNAH9* mutations. In the distal cilium of individuals with *DNAH9* mutations, we have speculatively left in a lightly shaded remnant ODA structure in the distal axoneme, since occasional outer dynein arms remain visible in TEM and electron tomography images and we cannot fully exclude their absence by immunofluorescence.

**Figure 4. Consistent location of the outer dynein arm gamma (DNAH5) and beta heavy chains (DNAH9 and DNAH11) of the ciliated epithelial cells at various nasal and bronchopulmonary sites.**

(A-C) Respiratory epithelial cells from a non-PCD donor were stained with a marker of the ciliary axoneme (acetylated alpha tubulin, green), the outer dynein arm beta and gamma heavy chain markers (DNAH9, DNAH11 or DNAH5, red) and the nucleus (DAPI, blue). (A) DNAH5 localises along the entire length of the cilia. (B) DNAH9 localises distally in the cilia and (C) DNAH11 localises proximally in the cilia (note that this antibody is associated with some non-specific background staining in the cytoplasm). The same pattern of staining localisation is seen for all three heavy chain proteins in epithelial cells dissected from both the peripheral airways of less than 1.5mm and from the lobe region, in comparison to the nasal epithelial cells shown in Figure 2. Scale bars, 5  $\mu$ M. (D) Cartoon showing the unified nasal and bronchopulmonary distribution of type 1 (DNAH5 and DNAH11-positive) and type 2 (DNAH5 and DNAH9-positive) ODAs in cilia throughout the airways.

**Figure 5. RNAi silencing of *Paramecium* DNAH9 shows severe defective motile cilia phenotype**

(A) Quantitative RT-PCR assessment of expression level of *DNAH9* orthologues after RNAi knockdown in *Paramecium*. The expression level is reduced by > 80% in the *DNAH9*-RNAi cells compared to its expression in the control *ND7*-RNAi cells (*ND7* gene not associated with cilia motility);  $p < 0.001$ , independent samples t test. (B) Reduced cilia beat frequency after *DNAH9*-RNAi silencing. At 48 hr of RNAi the ciliary beat frequency of *DNAH9* knockdown cells is reduced by ~50% compared to *ND7* control knockdown cells. 7–10 paramecia evaluated per condition;  $p < 0.001$ , Mann-Whitney U test. (C) Analysis of swimming velocity of *Paramecium* *DNAH9* and *ND7* knockdown cells shows a significant reduction (~70%) in velocity of *DNAH9* knockdown cells compared to the control *ND7* knockdown cells over 72 hours of RNAi silencing. More than 150 *Paramecia* were evaluated per condition;  $p < 0.001$ , independent samples t test. (D) *Paramecium* swimming pattern visualised in dark-field microscopy, shown by the Z projection of track recordings, captured using a 10x objective after 24 and 48 hr of RNAi feeding. *DNAH9* knockdown cells (right) show a severe motility phenotype compared to the control *ND7*

silenced cells (left). (E) Transmission electron micrographs of *Paramecium* cilia in cross-section showing normal 9+2 arrangement for the control *ND7* knockdown (left, outer dynein arms arrowed) and absence of the outer (grey arrows) dynein arms in the *DNAH9* knockdown cells (right). (F) Quantification of TEM dynein arms counts across >40 cross-sections per knockdown experiment showed reduced ODA and IDA in *DNAH9* knockdown cells compared to *ND7* control knockdown cells with ODA loss more extensive than IDA loss.  $p > 0.001$ , independent samples t test. All error bars indicate SEM.

**Table 1. Clinical features of individuals carrying *DNAH9* mutations**

<b>Individual:</b>	<b>1.II.1</b>	<b>2.II.2</b>	<b>2.II.3</b>	<b>3.II.1</b>
<i>DNAH9</i> variants NM_001372.3	c.12367G>A; p.Asp4123Asn + c.12367G>A; p.Asp4123Asn	c.8708-2A>G; p.Glu2904Aspfs*53 + c.10193G>T; p.Arg3398Leu	c.8708-2A>G; p.Glu2904Aspfs*53 + c.10193G>T; p.Arg3398Leu	c.5641A>G; p. Lys1881Glu + c.5641A>G; p. Lys1881Glu & c.8894G>A; p.Arg2965His + c.8894G>A; p.Arg2965His
Age at diagnosis	8 yrs	2 yrs	9 yrs	13 yrs
Neonatal distress	N	N	N	N
Rhinosinusitis	Y	Y	Y	Y
Wet cough	Y	Y with infection	Y with infection	Y
Otitis media	N	N	N	N
Bronchiectasis	N	N	N	N
Situs Inversus	Y	Y	Y	Y
Infertility	NA too young	NA too young	NA too young	Y
Lung function - forced expiratory volume in 1 second (FEV1%pred)	49	108	normal range	92
Nasal nitric oxide (normal >77nl/min)	105nl/min	46nl/min	100nl/min	216nl/min
Ciliary beat frequency (normal 8-15 Hz)	8.4-10.1Hz	8-9.3Hz	8.4-8.6Hz	5.0-6.6Hz
Co-morbidities	Protein losing enteropathy, complex congenital heart disease	N	N	N